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Description
      2384033
                DRUG? ?
S1
                ANTIVIRAL? ?
        45592
S2
       100722
                 SUSCEPTIBILITY
S3
                 RESISTANCE
S4
      1519592
                HEPATITIS (W) C (W) VIRUS
         9768
S5
         9688
                HCV
S 6
                 ANTI(W) VIRAL? ?
s7
         5591
                 RESISTANCE (W) TEST (W) VECTOR
S8
       306800
                 INDICATOR OR REPORTER
S 9
      1077419
                 EXPRESS?
S10
                 S1 OR S2 OR S7
      2405761
S11
      1587587
                 S3 OR S4
S12
                 S5 OR S6
S13
        13148
          346 . INTERNAL (W) RIBOSOME (W) ENTRY (W) SITE
S14
          676
                 IRES
S15
                 S14 OR S15
S16
          812
                 S11(S)S12(S)S13
S17
          191
                 VECTOR? ? OR PLASMID? ? OR COSMID? ?
       302192
S18
                 S17 AND S18
S19
           136
           133
                 S19 AND S10
S20
                 S20 NOT PY>1997
           38
S21
                 RD (unique items)
           38
S22
            66
                 S17 NOT PY>1997
S23
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S24
                 S11/AB
       408224
S25
       757126
                 S12/AB
S26
                 S13/AB
S27
          9385
                 S25 AND S26
S28
         26463
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                 S28 AND S13
S29
            15
                 S29 NOT PY>1997
S30
            15
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S31
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S32
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               OR AU="CAPON DANIEL JEFFREY US" OR AU="CAPON DANIEL JEFFRY" -
              OR AU="CAPON DJ"
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              JM"
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            21
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S36
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S41
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             6
S42
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S43
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S44
                 S13 AND S11 AND S12
           348
S45
S46
           157
                  S45 NOT S17
            74
                  S46 NOT PY>1997
S47
            74
                  RD (unique items)
S48
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8/3/1 (Item 1 from file: 340)
DIALOG(R) File 340:CLAIMS(R)/US Patent
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3072560 9839091

C/COMPOSITIONS AND METHODS FOR DETERMINING ANTI-VIRAL DRUG SUSCEPTIBILITY AND RESISTANCE AND ANTI-VIRAL DRUG SCREENING; MEASUREMENT, CALIBRATION GENE EXPRESSION

Inventors: Capon Daniel (US); Petropoulos Christos J (US)

Assignee: ViroLogic Inc Assignee Code: 47675

Applic Applic Issue Patent Number Date Date Number _____ _____ US 5837464 19981117 US 97790963 19970129 Patent: 19970129 US 97790963 Priority Applic: US 60-10715 19960129 Provisional Applic:

Calculated Expiration: 20170129

8/3/2 (Item 1 from file: 654)

DIALOG(R) File 654:US Pat. Full.

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02874739

Utility

COMPOSITIONS AND METHODS FOR DETERMINING ANTI-VIRAL DRUG SUSCEPTIBILITY AND RESISTANCE AND ANTI-VIRAL DRUG SCREENING [Measurement, calibration gene expression]

PATENT NO.: 5,837,464

ISSUED: November 17, 1998 (19981117)

INVENTOR(s): Capon, Daniel, Hillsborough, CA (California), US (United

States of America)

Petropoulos, Christos J., Half Moon Bay, CA (California), US

(United States of America)

ASSIGNEE(s): ViroLogic, Inc , (A U.S. Company or Corporation), S. San

Francisco, CA (California), US (United States of America)

[Assignee Code(s): 47675]

APPL. NO.: 8-790,963

FILED: January 29, 1997 (19970129)

FULL TEXT: 6433 lines

?

22/3,AB/1 (Item 1 from file: 654)

DIALOG(R) File 654:US Pat. Full.

(c) format only 1999 The Dialog Corp. All rts. reserv.

02724627

Utility

METHODS AND COMPOSITIONS FOR INHIBITING PRODUCTION OF REPLICATION COMPETENT VIRUS

[**Vector** directing **expression** of retroviral structural polypeptide comprising promoter, gene construct comprising nucleic acid molecule coding for polypeptide and biologically inactive inhibitory molecule, polyadenylation signal]

PATENT NO.: 5,698,446

ISSUED: December 16, 1997 (19971216)

INVENTOR(s): Klump, Wolfgang M., Del Mar, CA (California), US (United

States of America)

Jolly, Douglas J., Leucadia, CA (California), US (United

States of America)

ASSIGNEE(s): Chiron Corporation, (A U.S. Company or Corporation), US

(United States of America)
[Assignee Code(s): 11661]

APPL. NO.: 8-305,699

FILED: September 07, 1994 (19940907)

FULL TEXT: 2013 lines

ABSTRACT

The present invention provides methods and compositions for inhibiting the production of replication competent virus. The invention comprises nucleic acid cassettes encoding a non-biologically active inhibitory molecule which are incorporated into packaging cells and recombinant vector constructs. Upon recombination between various vector construct contained within the producer cell, a biologically active molecule is produced which kills the cell, thereby inhibiting production of replication competent virus.

22/3,AB/6 (Item 6 from file: 654)

DIALOG(R) File 654:US Pat. Full.

(c) format only 1999 The Dialog Corp. All rts. reserv.

02700749

Utility

RIBONUCLEASE RESISTANT VIRAL RNA STANDARDS

[Nucleic acid encapsulated by bacteriophage protein]

PATENT NO.: 5,677,124

ISSUED: October 14, 1997 (19971014)

INVENTOR(s): DuBois, Dwight B., Austin, TX (Texas), US (United States of

America)

Winkler, Matthew M., Austin, TX (Texas), US (United States of

America)

Pasloske, Brittan L., Austin, TX (Texas), US (United States of

America)

ASSIGNEE(s): Ambion, Inc , (A U.S. Company or Corporation), Austin, TX

(Texas), US (United States of America)

Cenetron Diagnostics LLC, (A U.S. Company or Corporation),

Austin, TX (Texas), US (United States of America)

[Assignee Code(s): 32084; 43403]

APPL. NO.: 8-675,153

FILED: July 03, 1996 (19960703)

FULL TEXT: 1847 lines

ABSTRACT

The present invention is directed to the process of creating a recombinant nucleic acid standard which is resistant to ribonuclease digestion and is non-infectious. A single strand of recombinant nucleic acid is encapsidated by bacteriophage proteins. The recombinant nucleic acid is a hybrid sequence encoding bacteriophage proteins and a specific non-bacteriophage sequence. A non-bacteriophage RNA sequence can be used as an RNA standard to help quantify the number of RNA molecules in an unknown sample. The recombinant RNA in its packaged form is highly resistant to ribonucleases, insuring that the RNA standard is not compromised by inadvertent ribonuclease contamination. These "ARMORED RNA" standards are ideal as RNA standards for the quantification of RNA viruses such as HIV and HCV from human body fluids such as blood and cerebrospinal fluid.

22/3,AB/7 (Item 7 from file: 654)

DIALOG(R) File 654:US Pat. Full.

(c) format only 1999 The Dialog Corp. All rts. reserv.

02693166

Utility

RECOMBINANT FOWLPOX VIRUS

PATENT NO.: 5,670,367

ISSUED: September 23, 1997 (19970923)

INVENTOR(s): Dorner, Friedrich, Vienna, AT (Austria)

Scheiflinger, Friedrich, Orth/Donau, AT (Austria) Falkner, Falko Gunter, Mannsdorf, AT (Austria)

ASSIGNEE(s): Immuno Aktiengesellschaft, (A Non-U.S. Company or Corporation)

, Vienna, AT (Austria)
[Assignee Code(s): 53908]

APPL. NO.: 8-232,463

FILED: April 22, 1994 (19940422)

PRIORITY: 91114300, EP (European Patent Office), August 26, 1991

(19910826)

This application is a continuation of application Ser. No. 07-935,313, filed Aug. 26, 1992, abandoned.

FULL TEXT: 4665 lines

ABSTRACT

An improved method is described to prepare recombinant fowlpox virus for the expression of proteins or for use as a vaccine. The new method uses for the insertion of foreign DNA an intergenic region which is located between the FPV thymidine kinase (tk)gene and the 3'-open reading frame. Said intergenic region is enlarged to comprise one or more unique restriction sites, thereby allowing insertion of foreign DNA in such a way that the FPV tk-gene remains intact and codes for the entire thymidine kinase. New strong poxvirus promoters are presented and new FPV host virus strains carrying a vaccinia virus thymidine kinase gene and an E. coli lacZ gene as a novel non-essential site. The novel fowlpox virus host strains allow the use of any insertion plasmid carrying vaccinia virus tk-gene flanking regions.

22/3,AB/22 (Item 22 from file: 654)

DIALOG(R) File 654:US Pat. Full.

(c) format only 1999 The Dialog Corp. All rts. reserv.

02612413

Utility HEPATITIS B VIRUS MUTANTS, REAGENTS AND METHODS FOR DETECTION [Diagnosis, vaccines]

PATENT NO.: 5,595,739

January 21, 1997 (19970121) ISSUED:

INVENTOR(s): Carman, William F., Glasgow, GB (United Kingdom)

Decker, Richard H., Deerfield, IL (Illinois), US (United

States of America)

Wallace, Lesley, Glasgow, GB (United Kingdom)

Mimms, Larry T., Lake Villa, IL (Illinois), US (United States

of America)

Solomon, Larry R., Mundelein, IL (Illinois), US (United States

of America)

ASSIGNEE(s): Abbott Laboratories, (A U.S. Company or Corporation), Abbott

Park, IL (Illinois), US (United States of America)

[Assignee Code(s): 152]

APPL. NO.: 8-59,031 FILED: May 07, 1993 (19930507) FULL TEXT: 2059 lines

ABSTRACT

Mutant Hepatitis B Virus (HBV) nucleic acid sequences useful for a variety of diagnostic and therapeutic applications, kits for using the HBV nucleic acid sequences, HBV immunogenic particles, and a method for producing antibodies to HBV. Also provided are methods for producing antibodies, polyclonal or monoclonal, from the HBV nucleic acid sequences.

(Item 1 from file: 155) 31/3,AB/1

DIALOG(R) File 155:MEDLINE(R)

(c) format only 1999 Dialog Corporation. All rts. reserv.

09690467 97449199

Therapy of hepatitis C: overview.

Lindsay KL

Department of Medicine, University of Southern California, Los Angeles 90033-4581, USA.

Sep 1997, 26 (3 Suppl 1) p715-77S, ISSN Hepatology (UNITED STATES) 0270-9139 Journal Code: GBZ

Languages: ENGLISH

Document type: CONSENSUS DEVELOPMENT CONFERENCE; CONSENSUS DEVELOPMENT CONFERENCE, NIH; JOURNAL ARTICLE; REVIEW

Based on the first decade of research on alpha interferon in viral hepatitis, one can conclude that up to 40% of patients with compensated chronic hepatitis C and elevated alanine aminotransferase (ALT) levels will respond at least transiently to interferon. Four forms of alpha interferon have been evaluated in large numbers of patients with chronic hepatitis C: alfa-2b, alfa-2a, alfa-n1, and consensus interferon (CIFN). Responses are defined on the basis of biochemical (ALT) or virological (hepatitis virus [HCV] RNA testing by polymerase chain reaction [PCR]) end points, as end-of-treatment response (ETR) or sustained response (SR). Biochemical ETR rates to 6 months of therapy range from 35% to 50%, and SR rates 6 months after treatment from 8% to 21%. Although 6-month treatment courses are associated with a significant rate of relapse, 12 months of initial treatment and re-treatment regimens markedly improve the SR rate. Long-term follow-up evaluation in patients with an SR to interferon consistently show long-lasting and significant clinical, virological, and histological improvement. Finally, baseline factors that have been shown to be associated with SR to 6 months of treatment are not accurate enough to predict response. Therefore, the best treatment strategy is a therapeutic trial. Further studies of interferon therapy of hepatitis C are needed to define better virological end points useful in stopping therapy, to understand and better manage significant side effects of interferon, and to of interferon in biochemical effects histological evaluate the nonresponders. Also needed is a better understanding of the causes of resistance to interferon. Finally, newer therapeutic regimens such as the use of induction therapy and combination therapies with ribavirin, other agents, cytokines, and cytokine modifiers are of primary antiviral importance in eventually developing safe and effective means of treatment of hepatitis C.

(Item 2 from file: 155) 31/3,AB/2

DIALOG(R) File 155:MEDLINE(R)

(c) format only 1999 Dialog Corporation. All rts. reserv.

09683658 97193758

Semiquantitative analysis of intrahepatic cytokine mRNAs in chronic hepatitis C.

Dumoulin FL; Bach A; Leifeld L; El-Bakri M; Fischer HP; Sauerbruch T;

Department of General Internal Medicine and Institute of Pathology, University of Bonn, Germany.

Mar 1997, 175 (3) p681-5, ISSN 0022-1899 J Infect Dis (UNITED STATES) Journal Code: IH3

Languages: ENGLISH

Document type: JOURNAL ARTICLE

In order to characterize intrahepatic cytokine production, the mRNA levels of interleukin (IL)-2, -4, and -10 and interferon (IFN)-gamma were semiquantitatively determined by reverse-transcription polymerase chain reaction in liver specimens from patients with chronic hepatitis C (n = 23), chronic hepatitis B (n = 9), or primary biliary cirrhosis (n = 12) and normal liver (control) specimens (n = 12). IL-4 mRNA was undetectable. Similar IL-10 mRNA levels were detected in all samples studied, including the controls. Mean IFN-gamma and IL-2 mRNA levels were elevated in chronic inflammatory liver disease. IL-2 mRNA levels were similar in all 3 patient groups, but intrahepatic IFN-gamma mRNA levels were significantly higher in chronic hepatitis C than in chronic hepatitis B or primary biliary cirrhosis patients. This predominance of IFN-gamma may indicate a lower susceptibility of hepatitis C virus to the antiviral effects of this cytokine. The presence of IL-10 in normal liver may impair the induction of antiviral immune responses.

31/3,AB/9 (Item 9 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

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08603990 97201548

Analysis of hepatitis C virus quasispecies populations by temperature gradient gel electrophoresis.

Lu M; Funsch B; Wiese M; Roggendorf M

Institut fur Virologie, Universitatsklinikum Essen, Germany.

J Gen Virol (ENGLAND) Apr 1995, 76 (Pt 4) p881-7, ISSN 0022-1317

Journal Code: I9B Languages: ENGLISH

Document type: JOURNAL ARTICLE

C virus (HCV) forms complex quasispecies populations Hepatitis which consist of a large number of closely related genetic variants. This genetic heterogeneity may cause antigenic variation or drug resistance . We used heteroduplex analysis by temperature gradient gel electrophoresis (TGGE) to characterize genetic variants of \mathbf{HCV} . The high resolution of TGGE was proven by comparison of DNA sequence data of different cDNA clones from the HCV 5'NCR with their corresponding migration pattern in TGGE. Using this method we were able to identify virus variants of the HCV 5'NCR even if they only differed from each other by a single base. HCV populations from three patients with chronic hepatitis C were found to consist of genetic variants, although the degree of the heterogeneity varied. In addition, we compared the genetic heterogeneity of the core and E2 regions of the HCV genome in one patient. Our results demonstrate that TGGE is a useful tool for characterization of the genetic heterogeneity of virus populations in vivo.

31/3,AB/10 (Item 10 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

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08591853 96311182

Clinical relevance of hepatitis C virus quasispecies.

Enomoto N; Sato C

Second Department of Internal medicine, Faculty of Medicine, Tokyo Medical and Dental University, Japan.

J Viral Hepat (ENGLAND) 1995, 2 (6) p267-72, ISSN 1352-0504

Journal Code: CG0

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

It has been shown that hepatitis C virus (HCV) populations in infected individuals are composed of quasispecies with diverse mutations. The analysis of these variants may reveal mechanisms of the persistence of HCV infection, carcinogenesis and resistance to antiviral therapy. Recently, genetic features of interferon-resistant HCV have been elucidated through the analysis of interferon-resistant quasispecies, making it possible to predict interferon efficacy by detecting interferon-resistant strains.

41/3,AB/1 (Item 1 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

(c) format only 1999 Dialog Corporation. All rts. reserv.

09296938 98007644

Efficient gene transfer into various mammalian cells, including non-hepatic cells, by baculovirus vectors.

Shoji I; Aizaki H; Tani H; Ishii K; Chiba T; Saito I; Miyamura T;

Department of Virology II, National Institute of Infectious Diseases, Tokyo, Japan.

J Gen Virol (ENGLAND) Oct 1997, 78 (Pt 10) p2657-64, ISSN 0022-1317 Journal Code: I9B

Languages: ENGLISH

Document type: JOURNAL ARTICLE

baculovirus (Autographa californica nucleopolyhedrovirus) vector containing a strong promoter, the CAG promoter, was developed to introduce foreign genes into mammalian cells. Recombinant baculoviruses carrying a reporter gene under the control of the CAG promoter were inoculated into various mammalian cell lines. High-level expression was observed not only in hepatocytes but also in other non-hepatic cell lines tested. Expression of the reporter gene was detected even 14 days after infection. The infectious titre of the recovered baculoviruses decreased significantly after infection, indicating that the baculoviruses did not replicate in mammalian cells. We then compared the efficiencies of gene expression by the baculovirus vector with that of a replication-defective adenovirus by using the same expression unit. The same level of expression was observed in HepG2, HeLa and COS7 cells by both vectors . Efficient expression and proper processing were observed in mammalian cells infected baculoviruses carrying genes coding for structural regions of virus . These results suggest that the baculovirus vector C is a good tool for gene delivery into various mammalian cells in order to study the function of foreign genes.

41/3,AB/2 (Item 2 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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09276812 97415421

In vivo translational efficiency of different hepatitis C virus 5'-UTRs.

Buratti E; Gerotto M; Pontisso P; Alberti A; Tisminetzky SG; Baralle FE
International Centre for Genetic Engineering and Biotechnology, Trieste,

Italy.

FEBS Lett (NETHERLANDS) Jul 14 1997, 411 (2-3) p275-80, ISSN 0014-5793 Journal Code: EUH

Languages: ENGLISH

Document type: JOURNAL ARTICLE

Initiation of translation in hepatitis C virus (HCV) is dependent on the presence of an internal ribosome entry site (IRES) contained in its 341-nt-long 5'-untranslated region (UTR). This region is very conserved among different isolates and has been used to classify HCV isolates in six different genotypes. These genotypes differ in nucleotide sequence that generally preserve the IRES structure. However, the small differences seen may be biologically and clinically significant as the HCV strains seem to differ from each other in several important ways, such as the behaviour of the viral infection and the response to interferon therapy. Therefore, differences in translational initiation efficiency amongst the various genotypes could provide an explanation for these phenomena. Using a bicistronic expression system we have compared the in vivo translational ability of the three most common European genotypes of HCV (1, 2, and 3). The results show that genotype 3 is less able than 1 and 2 to promote translation initiation. In addition, by site-directed mutagenesis of the sequence of the IRES domain III apical stem loop structure, we have shown that the conservation of the primary nucleotide sequence and not only the structure, is important for the conservation of IRES function.

41/3,AB/7 (Item 7 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

(c) format only 1999 Dialog Corporation. All rts. reserv.

08260672 95133223

Pestivirus translation initiation occurs by internal ribosome entry.

Poole TL; Wang C; Popp RA; Potgieter LN; Siddiqui A; Collett MS

Oak Ridge National Laboratory, Biology Division, Tennessee 37831.

Virology (UNITED STATES) Jan 10 1995, 206 (1) p750-4, ISSN 0042-6822 Journal Code: XEA

Contract/Grant No.: HL43375, HL, NHLBI

Languages: ENGLISH

Document type: JOURNAL ARTICLE

The role of the 385 nucleotide 5' noncoding region (NCR) in the translation of the pestivirus genome was investigated. In vitro translation of an RNA transcript containing the 5' NCR of the bovine viral diarrhea virus (BVDV) genome followed by the coding sequence of the first gene product (p20) of the BVDV large open reading frame resulted in the synthesis of a 20-kDa polypeptide. Results from hybrid-arrest translation studies identified a region involving a predicted RNA stem-loop structure spanning nucleotides 154-261 within the 5' NCR that was important for p20 synthesis. An additional inhibitory oligonucleotide was complementary to the sequence at the base of this stem-loop and encompassed the initiating AUG at nucleotide 386. Antisense oligonucleotides both upstream and downstream of those that were inhibitory had no effect on p20 translation. RNA from a dicistronic expression vector in which the BVDV 5' NCR was inserted between two reporter genes, CAT and LUC, showed strong expression of the second (LUC) cistron upon in vitro translation. This expression was dramatically reduced in an analogous construct in which nucleotides 173-236 of the 5' NCR were deleted. Similar results were obtained when RNA from these same vectors was evaluated for expression after transfection into BHK cells. These results suggest that the BVDV 5' NCR contains an internal ribosome entry site for translation initiation. This translational mechanism is similar to that shown for hepatitis virus , further demonstrating the close relationship between viruses of these two genera within the family Flaviviridae.

41/3,AB/12 (Item 2 from file: 654)

DIALOG(R) File 654:US Pat. Full.

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02690507

Utility

MAMMALIAN EXPRESSION SYSTEMS FOR HCV PROTEINS

PATENT NO.: 5,667,992

ISSUED: September 16, 1997 (19970916)

INVENTOR(s): Casey, James M., Zion, IL (Illinois), US (United States of

America)

Bode, Suzanne L., Zion, IL (Illinois), US (United States of

America)

Zeck, Billy J., Gurnee, IL (Illinois), US (United States of

America)

Yamaguchi, Julie, Chicago, IL (Illinois), US (United States of

America)

Frail, Donald E., Libertyville, IL (Illinois), US (United

States of America)

Desai, Suresh M., Libertyville, IL (Illinois), US (United

States of America)

Devare, Sushil G., Northbrook, IL (Illinois), US (United

States of America)

ASSIGNEE(s): Abbott Laboratories, (A U.S. Company or Corporation), Abbott Park, IL (Illinois), US (United States of America)

[Assignee Code(s): 152]

APPL. NO.: 8-453,552

May 30, 1995 (19950530) FILED:

This is a division of U.S. patent application Ser. No. 08-417,478 filed Apr. 5, 1995 now abandoned, which is a continuation of 08-144,099 filed Oct. 28, 1993 now abandoned, which is a continuation of 07-830,024 filed Jan. 31, 1992, now abandoned.

7353 lines FULL TEXT:

ABSTRACT

Mammalian expression systems for the production of HCV proteins. Such expression systems provide high yields of HCV proteins, and enable the development of diagnostic and therapeutic reagents which contain glycosylated structural antigens and also allow for the isolation of the HCV etiological agent.

(Item 3 from file: 654) 41/3,AB/13

DIALOG(R) File 654:US Pat. Full.

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02627460

Utility

MAMMALIAN EXPRESSION SYSTEMS FOR HEPATITIS C VIRUS ENVELOPE GENES

PATENT NO.: 5,610,009

March 11, 1997 (19970311) ISSUED:

INVENTOR(s): Watanabe, Shinichi, Northbrook, IL (Illinois), US (United

States of America)

Yamaguchi, Julie, Chicago, IL (Illinois), US (United States of

America)

Desai, Suresh M., Libertyville, IL (Illinois), US (United

States of America)

Devare, Sushil G., Northbrook, IL (Illinois), US (United

States of America)

ASSIGNEE(s): Abbott Laboratories, (A U.S. Company or Corporation), Abbott

Park, IL (Illinois), US (United States of America)

[Assignee Code(s): 152]

8-188,281 APPL. NO.:

January 28, 1994 (19940128) FILED:

FULL TEXT: 2816 lines

ABSTRACT

Mammalian expression systems for the production of HCV E1-E2 fusion proteins. Such expression systems provide high yields of **HCV** proteins extracelluarly, and enable the development of diagnostic, vaccine and therapeutic reagents which contain glycosylated structural antigens and also allow for the isolation of the HCV etiological agent.

48/3,AB/1 (Item 1 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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09644116 98340243

Synthetic antisense oligodeoxynucleotides as potential drugs against hepatitis C.

Caselmann WH; Eisenhardt S; Alt M

Department of General Internal Medicine, Rheinische Friedrich-Wilhelms-Universitat, Bonn, Germany. Caselmann@Uni-Bonn.de Intervirology (SWITZERLAND) 1997, 40 (5-6) p394-9, ISSN 0300-5526 Journal Code: GW7

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

Antisense oligodeoxynucleotides (ODNS) can be used to specifically inhibit hepatitis C viral gene expression. Due to its high degree of conservation and its important function as internal ribosomal entry site, the 5'-non-coding region of the hepatitis C virus has been the most effective target to inhibit translation so far. Inhibition of luciferase reporter gene expression of up to 96 +/- 2% has been achieved. like phosphorothioate, methylphosphonate or ODNs Modifications of modification of terminal or intramolecular benzylphosphonate internucleotide phosphates lead to altered lipophilicity and binding stability to its RNA target and resistance against serum nucleases. The mode of action of ODNs is mainly dependent on an efficient induction of RNase H activity. The uptake of ODNs occurs via receptor-mediated or absorptive and fluid-phase endocytosis. After release from the endosomes, ODNs may exert their effects by interaction with cytosolic or nuclear structures. Side effects can occur when this interaction affects intra- or extracellular targets essential for biological cell function. If these problems can be solved, antisense technology has the potential for future therapy of human disease.

48/3,AB/3 (Item 3 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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09310408 98010017

Different susceptibilities of genetic variants of hepatitis C virus (HCV) to interferon (IFN).

Lu M; Wiese M; Funsch B; Roggendorf M

Institut fur Virologie, Universitatsklinikum Essen, Federal Republic of Germany.

Arch Virol (AUSTRIA) 1997, 142 (3) p581-8, ISSN 0304-8608 Journal Code: 8L7

Languages: ENGLISH

Document type: JOURNAL ARTICLE

Genetic variants of HCV may have different degrees of resistance to IFN and may therefore influence the outcome of IFN therapy. However, selection of HCV variants by IFN has not been investigated in detail. In this paper, heteroduplex analysis was used to monitor major changes of HCV populations in 4 chronically infected patients under IFN therapy. We found that a major variant of the HCV 5' non-coding region (5' NCR) emerged in a responder. In other patients although no new variant of the 5' NCR was identified, significant changes occurred within the core and El region of the HCV genome. Disappearance and emergence of HCV variants may reflect their different susceptibilities to IFN. Our results indicate that responses of HCV populations to IFN are complex and need to be characterized by analysis of multiple HCV genome regions.

48/3,AB/5 (Item 5 from file: 155)
DIALOG(R)File 155:MEDLINE(R)

(c) format only 1999 Dialog Corporation. All rts. reserv.

09296947 97475410

Mutations in the NS5A gene of hepatitis C virus in North American patients infected with HCV genotype 1a or 1b.

Hofgartner WT; Polyak SJ; Sullivan DG; Carithers RL Jr; Gretch DR Department of Laboratory Medicine, University of Washington Medical Center, Seattle, USA.

J Med Virol (UNITED STATES) Oct 1997, 53 (2) p118-26, ISSN 0146-6615 Journal Code: I9N

Contract/Grant No.: R29 AI39049-02, AI, NIAID; AI/DK 41320-02, AI, NIAID Languages: ENGLISH

Document type: JOURNAL ARTICLE

Previous studies from Japan have described an association between a conserved sequence within the hepatitis C virus (HCV) genome and resistance to interferon (IFN) therapy for patients infected with HCV genotype 1b [Enomoto et al. (1995): Journal of Clinical Investigation 96: 224-230; Enomoto et al. (1996): New England Journal of Medicine 334:77-81]. The present study examines amino acid sequences surrounding the putative Interferon Sensitivity Determining Region (ISDR) of the NS5A gene of HCV in 21 North American patients with genotype 1a or 1b infection receiving recombinant IFN therapy. The ISDR consensus or intermediate pattern was observed in 13 of 14 NS5A clones from North American patients infected with genotype 1b. However, we found no evidence of the consensus ISDR sequence in any NS5A clones isolated from 15 patients with genotype la infection. In select cases, gel shift analysis showed no significant changes in the clonal frequency of the putative ISDR domain of HCV genotype la or lb infected patients who were either nonresponsive to IFN therapy, or relapsed following withdrawal of IFN therapy. These results suggest that a conserved domain is neither associated with, nor responsible for, IFN resistance in North American patients infected with HCV genotype la, and demonstrate a need for further investigation into the reported association between ISDR consensus sequences and IFN resistance in genotype 1b clones.

48/3,AB/8 (Item 8 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

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09260019 97201443

Pretreatment virus load and multiple amino acid substitutions in the interferon sensitivity-determining region predict the outcome of interferon treatment in patients with chronic genotype 1b hepatitis C virus infection [see comments]

Chayama K; Tsubota A; Kobayashi M; Okamoto K; Hashimoto M; Miyano Y; Koike H; Kobayashi M; Koida I; Arase Y; Saitoh S; Suzuki Y; Murashima N; Ikeda K; Kumada H

Department of Gastroenterology, Toranomon Hospital, Minato-Ku, Tokyo, Japan.

Hepatology (UNITED STATES) Mar 1997, 25 (3) p745-9, ISSN 0270-9139 Journal Code: GBZ

Comment in Hepatology 1997 Mar; 25(3):769-71

Languages: ENGLISH

Document type: JOURNAL ARTICLE

Hepatitis C virus (HCV) genotype 1b and high pretreatment virus load are predictive factors of poor response to interferon therapy in patients with chronic hepatitis C. To further examine the factors predicting the response to interferon in patients with genotype 1b infection, we analyzed 110 consecutive patients with HCV who were treated with a total of 624 million units of lymphoblastoid interferon alfa. Thirty-six patients (33%) were responders, while the remaining 74 patients (67%) were nonresponders. Multivariate analysis showed that a high virus titer (assessed by serum core protein level, P = .0021) and the presence of

amino acid substitutions in the interferon more two sensitivity-determining region (ISDR) (P = .0036) correlated significantly with the response to interferon therapy. Because mutations analyzed by direct sequencing of polymerase chain reaction (PCR) products may reflect artifacts of direct sequencing, we further analyzed quasispecies of HCV in this region by cloning and sequencing. Although PCR-based analysis of responders with multiple amino acid substitutions in the ISDR showed the presence of a small amount of wild-type strain in their serum, the results obtained by direct sequencing and cloning were essentially the same. A longitudinal study of quasispecies in 2 patients who showed a dramatic change in the virus titer showed no conversion from wild type to mutant or vice versa. Our results indicate that amino acid substitutions and virus load are independent predictors of the response to interferon therapy. The ability of some patients with no mutation in the ISDR or high virus load to eliminate the virus suggests the presence of other unidentified factors, host or viral, that influence the response to interferon therapy.

48/3,AB/9 (Item 9 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

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09260018 97201442

Mutations in the nonstructural 5A gene of European hepatitis C virus isolates and response to interferon alfa [see comments]

Zeuzem S; Lee JH; Roth WK

Medizinische Klinik II, Klinikum der Johann Wolfgang Goethe-Universitat, Frankfurt a.M., Germany.

Hepatology (UNITED STATES) Mar 1997, 25 (3) p740-4, ISSN 0270-9139 Journal Code: GBZ

Comment in Hepatology 1997 Mar; 25(3):769-71

Languages: ENGLISH

Document type: JOURNAL ARTICLE

The response rate to interferon alfa (IFN-alpha) in patients infected with hepatitis C virus (HCV) genotype 1 isolates is poor. A region associated with sensitivity to IFN has been identified in subtype HCV -1b isolates from Japanese patients in the carboxyterminal half of the nonstructural protein NS5A (between codon 2209 and 2248). HCV -1b isolates with at least four amino acid changes in this region compared with the HCV -1b prototype sequence were sensitive, whereas isolates identical to the prototype sequence were resistant to IFN-alpha. Patients infected with HCV -1b isolates carrying 1 to 3 mutations in NS5A(2209-2248) showed an of the large geographical response pattern. Because intermediate differences observed for HCV it is unknown whether this putative IFN-alpha sensitivity determining region is also predictive for European isolates. We analyzed 32 patients chronically infected with HCV -la or HCV -1b isolates who were treated with 3 million units of recombinant IFN-alpha three times per week for 1 year. Before initiation, during, and after treatment serum HCV -RNA levels were assessed by a quantitative reverse-transcription polymerase chain reaction (RT-PCR) assay. The amino acid sequence of NS5A(2209-2248) was determined by direct sequencing of the PCR-amplified HCV genome and was compared with the reference sequence HCV -J. In patients chronically infected with subtype HCV -la or HCV -lb the initial or sustained response to IFN-alpha was not related to the number of amino acid substitutions in the NS5A(2209-2248) region. In addition, the number of amino acid changes in NS5A(2209-2248) was not related to pretreatment HCV -RNA serum levels. In three patients with a pronounced initial decline of HCV -RNA levels (>3 log) sequence analyses of NS5A(2209-2248) were performed before and after therapy. Compared with the pretreatment amino acid sequence the HCV isolates of these patients revealed more mutations in the NS5A(2209-2248) region after therapy. These findings from European patients indicate that the NS5A(2209-2248) region of does not represent a common interferon sensitivity determining region.

48/3,AB/19 (Item 19 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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08589239 96155132

Hepatitis C virus quasispecies populations during chronic hepatitis C infection.

Enomoto N; Sato C

Second Dept of Internal Medicine, Faculty of Medicine, Tokyo Medical and Dental University, Japan.

Trends Microbiol (ENGLAND) Nov 1995, 3 (11) p445-7, ISSN 0966-842X Journal Code: B1N

Languages: ENGLISH

Document type: JOURNAL ARTICLE; REVIEW; REVIEW, TUTORIAL

Hepatitis C virus populations in infected individuals consist of quasispecies with diverse mutations. These quasispecies have different biological properties, and the analysis of these variants has led to new interpretations of viral persistence, carcinogenesis and **resistance** to interferon therapy.

48/3,AB/20 (Item 20 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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08587326 96091314

Predictors of sustained response, relapse and no response in patients with chronic hepatitis C treated with interferon-alpha.

Chemello L; Cavalletto L; Noventa F; Bonetti P; Casarin C; Bernardinello E; Pontisso P; Donada C; Casarin P; Belussi F; et al

Clinica Medica 2, University of Padova, PD, Italy.

J Viral Hepat (ENGLAND) 1995, 2 (2) p91-6, ISSN 1352-0504

Journal Code: CG0

Languages: ENGLISH

Document type: CLINICAL TRIAL; JOURNAL ARTICLE; RANDOMIZED CONTROLLED

of response are seen when interferon-alpha patterns main Three (IFN-alpha) is used for the treatment of chronic hepatitis C: 1 sustained response with alanine-aminotransferase (ALT) normalization that is maintained after cessation of therapy, with or without clearance of serum (HCV) RNA; 2 transient response with ALT C hepatitis virus normalization during therapy followed by relapse after its withdrawal, and 3 no response with no or only partial reduction in ALT levels. In order to define variables that could predict each of these three types of response we studied 321 cases of chronic hepatitis C treated with IFN-alpha in two consecutive trials conducted in our Unit. By univariate analysis, age < 45 years (P < 0.01), known disease duration < 60 months (P < 0.01), normal gamma-glutamyl-transpeptidase (gamma GT) levels (P < 0.01) and infection by genotype 2 or HCV genotype 3 (P < 0.01) were found to be statistically associated with sustained response while age > 45 years (P <0.01), body weight (P = 0.05), cirrhosis (P < 0.01) and elevated gamma GT (P < 0.01) were associated with no response. By multivariate analysis sustained response was predicted by HCV genotype 2 (P < 0.01) and HCV genotype 3 (P < 0.01), known disease duration (P < 0.01), patient's age (P < 0.05) and associated with the use of a more aggressive treatment schedule (P < 0.05). Transient response with relapse was predicted by known duration of disease (P < 0.05), HCV genotype 1 (P < 0.05) and female sex (P < 0.05). No response was statistically associated with elevated gamma GT levels (P < 0.01), higher body weight (P < 0.05) and with the less aggressive regimen of 3 MU of natural IFN-alpha given three times weekly for 6 months (P < 0.05). These results indicate that the HCV genotype as well as the schedule of treatment greatly affect the pattern of response to IFN in chronic hepatitis C and allow us to define criteria to predict which type of response is more likely in individual patients.

48/3,AB/23 (Item 23 from file: 155)

DIALOG(R) File 155: MEDLINE(R)

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08580799 95340824

Comparison of full-length sequences of interferon-sensitive and resistant hepatitis C virus 1b. Sensitivity to interferon is conferred by amino acid substitutions in the NS5A region.

Enomoto N; Sakuma I; Asahina Y; Kurosaki M; Murakami T; Yamamoto C; Izumi N; Marumo F; Sato C

Second Department of Internal Medicine, Faculty of Medicine, Tokyo Medical and Dental University, Japan.

J Clin Invest (UNITED STATES) Jul 1995, 96 (1) p224-30, ISSN 0021-9738 Journal Code: HS7

Languages: ENGLISH

Document type: JOURNAL ARTICLE

We have previously demonstrated that sensitivity to interferon is different among hepatitis C virus (HCV) quasispecies simultaneously in same individuals and that interferon-resistant HCV quasispecies are selected during the treatment. To determine the genetic basis of their resistance to interferon, HCV genotype-1b was obtained from serum of three patients before and during interferon therapy, and their full-length nucleotide and deduced amino acid sequences were determined. Comparison of the pairs of interferon-resistant and interferon-sensitive HCV isolates in respective individuals demonstrated clusters of amino acid differences in the COOH-terminal half of the NS5A region (codon 2154-2383), which contained a common unique amino acid difference at codon 2218. Additional sequence data of the COOH-terminal half of the NS5A region obtained from six interferon-resistant and nine interferon-sensitive HCV confirmed the exclusive existence of missense mutations in a 40 amino acid stretch of the NS5A region around codon 2218 (from codon 2209 to 2248) in interferon-sensitive HCV . On the other hand, this region of interferon-resistant HCV was identical to that of prototype HCV genotype-1b (HCV -J, HCV -JTa, or HC-J4). We designated this region as the interferon sensitivity determining region. Thus, HCV genotype-1b with the prototype interferon sensitivity determining region appears to be interferon-resistant strains. The specific nature of these mutations might make it possible to predict prognostic effects of interferon treatment.

48/3,AB/31 (Item 31 from file: 155)

DIALOG(R) File 155:MEDLINE(R)

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07693335 94069717

Increased susceptibility for CsA-induced hepatotoxicity in kidney graft recipients with chronic viral hepatitis C.

Horina JH; Wirnsberger GH; Kenner L; Holzer H; Krejs GJ

Department of Medicine, Karl Franzens University, Graz, Austria.

Transplantation (UNITED STATES) Nov 1993, 56 (5) p1091-4, ISSN 0041-1337 Journal Code: WEJ

Languages: ENGLISH

Document type: JOURNAL ARTICLE

CsA-induced hepatotoxicity is a rare disorder in renal transplant recipients when low doses are administered and whole blood trough levels of CsA are regularly monitored. However, there is controversy about the clinical value of measuring CsA-metabolites, whose contribution to immunosuppression and toxicity is not fully understood. To assess the relation between low-dose CsA therapy and hepatotoxicity, we studied 128

renal transplant recipients attending our nephrology clinic. Eight of these patients had markedly elevated liver function tests. Three patients while receiving very low doses of oral CsA (< 3.8 mg/kg of body weight) presented marked derangements of CsA metabolism with abnormally increased CsA-metabolite levels. Parent drug levels were in the normal range. All 3 patients had chronic infection with hepatitis C virus and revealed histomorphologic evidence of hepatotoxicity. Hepatic dysfunction normalized when CsA was withdrawn or reduced by 50%. It is likely that hepatitis C virus infection interferes with CsA metabolism and/or biliary CsA-excretion and thus is responsible for CsA and/or metabolite-induced hepatotoxicity despite very low doses of CsA.

(Item 6 from file: 348) 48/3,AB/40 DIALOG(R) File 348: European Patents (c) 1999 European Patent Office. All rts. reserv. 00765862 ORDER fax of complete patent from Dialog SourceOne. See HELP ORDER 348 Methods and compositions for controlling translation of HCV proteins Verfahren und Zusammensetzungen zur Kontrolle der Uberstetzung von HCV -Proteine Procedes et compositions pour le controle de la traduction des proteines de HCV PATENT ASSIGNEE: CHIRON CORPORATION, (572531), 4560 Horton Street, R440, Emeryville California 94608-2916, (US), (applicant designated states: AT; BE; CH; DE; DK; ES; FR; GB; GR; IE; IT; LI; LU; MC; NL; PT; SE) INVENTOR: Hang, Jan H., 3238 Del Mar Drive, Lafayette, CA 94549, (US) Spaete, Richard R., 2501 Cornonet Boulevard, Belmont, CA 94002, (US) Yoo, Byoung J., Han-Dong University, 3 Namsong-ri, Hunghae-eub, Puk-ku Puhnag, Kyungbuk, (KR) Suh, Byung S., Han-Dong University, 3 Namsong-ri, Hunghae-eub, Puk-ku Puhnag, Kyungbuk, (KR) Selby, Mark J., 136 Galewood Circle, San Francisco, CA 94131, (US) Houghton, Michael, 53 Rosemead Court, Danville, CA 94529, (US) LEGAL REPRESENTATIVE: Goldin, Douglas Michael (31061), J.A. KEMP & CO. 14 South Square Gray's Inn, London WC1R 5LX, (GB) PATENT (CC, No, Kind, Date): EP 718400 A2 960626 (Basic) EP 718400 A3 960703 EP 95118443 930928; APPLICATION (CC, No, Date): PRIORITY (CC, No, Date): US 952799 920928 DESIGNATED STATES: AT; BE; CH; DE; DK; ES; FR; GB; GR; IE; IT; LI; LU; MC; NL; PT; SE RELATED PARENT NUMBER(S) - PN (AN): EP 662128 (EP 939224143) INTERNATIONAL PATENT CLASS: C12N-015/11; C12N-015/51; C12N-015/67; C12N-015/86; A61K-031/70; A61K-047/48; A61K-048/00 ABSTRACT EP 718400 A3 Embodiments of the present invention feature methods and compositions for controlling the translation of viral peptides and proteins from viral nucleic acid, with particular applications to pestivirus and HCV . The methods and compositions feature control elements of the 5'UT region of the viral genome. ABSTRACT WORD COUNT: 54

LANGUAGE (Publication, Procedural, Application): English; English; Fulltext AVAILABILITY:
Available Text Language Update Word Count

Available Text Language Update Word Count CLAIMS A (English) EPAB96 336
SPEC A (English) EPAB96 8437

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Total word count - document A
Total word count - document B
                                      8773
Total word count - documents A + B
                (Item 11 from file: 348)
 48/3,AB/45
DIALOG(R) File 348: European Patents
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00562000
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RNA and DNA molecules for producing virus resistance.
DNS- und RNS-Molekule zur Erzeugung einer Virus-Resistenz.
Molecules d'ADN et d'ARN pour la production d'une resistance virale.
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PATENT (CC, No, Kind, Date): EP 558944 A2 930908 (Basic)
                              EP 558944 A3 940608
                              EP 93101710 930204;
APPLICATION (CC, No, Date):
PRIORITY (CC, No, Date): DE 4203441 920206
DESIGNATED STATES: BE; CH; DE; DK; FR; GB; IT; LI; NL
INTERNATIONAL PATENT CLASS: C12N-015/11; C12N-001/21; A01H-005/00;
  A01H-005/10; C12Q-001/68; A01N-063/02; C12N-009/00;
ABSTRACT EP 558944 A2
    The invention relates to RNA molecules which are complementary to at
  least one part of a viral RNA replicative intermediate and which inhibit
  the viral growth cycle by binding to the RNA replicative intermediate and
  preferably by specifically cleaving the RNA replicative intermediate,
  thereby achieving a reliable improvement in virus resistance in the
  desired organisms. (see image in original document)
ABSTRACT WORD COUNT: 62
LANGUAGE (Publication, Procedural, Application): English; English; English
FULLTEXT AVAILABILITY:
Available Text Language
                           Update
                                     Word Count
                                       586
      CLAIMS A (English)
                           EPABF1
               (English) EPABF1
                                      6493
      SPEC A
                                      7079
Total word count - document A
Total word count - document B
Total word count - documents A + B
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